Office européen des brevets

11) Publication number:

0 109 561

A1

Party

12

### **EUROPEAN PATENT APPLICATION**

21 Application number: 83110477.3

(6) Int. Cl.3: A 61 K 31/17

22 Date of filing: 20.10.83

(30) Priority: 27.10.82 US 437072

(43) Date of publication of application: 30.05.84 Bulletin 84/22

Designated Contracting States;
 AT BE CH DE FR GB IT LI LU NL SE

7 Applicant: USV PHARMACEUTICAL CORPORATION
1 Scarsdale Road
Tuckahoe New York(US)

(72) Inventor: Wolf, Peter S. Richard Somers Road Granite Springs New York(US)

(2) Inventor: Kasseiman, Morris A. 21 Autumn Circle Yonkers New York(US)

(74) Representative: Patentanwälte Grünecker, Dr. Kinkeldey, Dr. Stockmalr, Dr. Schumann, Jakob, Dr. Bezold, Meister, Hilgers, Dr. Meyer-Plath Maximilianstrasse 58 D-8000 München 22(DE)

64 Celiproiol for the treatment of glaucoma.

(5) An ophthalmic preparation for the treatment of glaucoma by the application to the glaucomatous eye of a celiprolol salt in a pharmaceutically acceptable ophthalmic carrier.

# CELIPROLOL FOR THE TREATMENT OF GLAUCOMA

This invention relates to compositions and methods for treating intraocular pressure associated with glaucoma. More particularly it relates to the use of celiprolol hydrochloride, as well as selected pharmaceutically acceptable salts thereof, that have been found useful in lowering intraocular pressure.

Elevated intraocular pressure is a major risk factor in the onset and development of glaucomatous visual field loss. The higher the level of intraocular pressure, the greater the likelihood of glaucomatous visual field loss and optic nerve damage.

Attempts have been made to lower intraocular pressure in glaucoma by administering the patients certain beta adrenergic blocking agents, commonly known as beta blockers.

In the human body, the beta blockers have several effects. For example, they can reduce the heart rate of an angina patient, which in turn reduces the workload of the heart and thus its need for blood and oxygen. They also tend to decrease the heart's force of contraction, which likewise diminishes the heart's workload. In addition, these drugs reduce the systolic blood pressure which is beneficial to patients with hypertension.

In addition to treating heart ailments, some beta blockers were experimented with for treating other ailments, such as migraine, alcohol and drug withdrawal problems, and glaucoma.

While the use of beta blockers provides valuable

benefits to humankind, it also has some undesirable side
effects and reactions. Some beta blockers tend to build up
in the central nervous system causing fatigue, lethargy, and

1

10

15

confusion. Others may cause bronchial spasm and cannot be used in people with bronchial asthma. Still others, while having minimal side effects, fail to produce acceptable results because of their limited potency.

Timolol, a non-selective beta-adrenoceptor 5 antagonist, has been shown to lower intraocular pressure in both patients with normal intraocular pressure and in patients with open angle glaucoma. Sold as TIMOPTIC (R) it is the only beta block sold in the U.S. for this purpose. Among the adverse reactions reported on the use of timolol is 10 the aggravation or precipitation of certain cardiovascular and pulmonary disorders including bronchospastic disease, sinus bradycardia, cardiogenic shock and cardiac failure. Similarly, atenolol, sotalol, pindolol, oxprenolol, practolol, propranolol, butidrine and metoprolol were reported to have activity in the treatment of glaucoma. However, some of these compounds have been reported to cause pronounced side~effects, for example, metoprolol provokes allergic reactions, exprendlol may cause corneal epitheliopathy and practolol induces oculomucoculaneous syndrome by immunopathological reactions.

Celiprolol hydrochloride has been shown a selective beta-1-adrenoceptor antagonist having intrinsic sympathomimetic, but no local anesthetic activity.

25. It has now been discovered that celiprolol hydrochloride, as well as selected pharmaceutically acceptable salts thereof such as maleate, succinate and the like, when topically applied to the eyes in a pharmaceutically suitable vehicle, such as an ophthalmic solution, is effective in lowering intraocular pressure.

Such ophthalmic preparation contain from about 0.01% w/v to about 5.0% w/v, preferably from about 0.03 w/v to about 2.0%

w/v of the active ingredient along with inactive ingredients used in the art, such as sodium borate-boric acid, sodium hydroxide to adjust pH, benzalkonium chloride as preservative and water as the vehicle of preference.

It has also been surprisingly discovered that celiprolol hydrochloride, when used according to the present invention, produces bronchodilation following each application and as such possesses significant therapeutic advantage to patients suffering from both glaucoma and respiratory disease.

Celiprolol hydrochloride (3-[3-acetyl-4-[3(tert-butylamino)-2-hydroxy-propoxy]-phenyl]-1, 1-diethylurea hydrochloride) as the free base has the following structure:

It has a molecular weight of 415.95 having C =

57.75%, H = 8.24%, N = 10.10%, 0 = 15.39% and Cl = 8.52%.

Celiprolol can be prepared according to several pathways described in Austrian Patents: 334,385 issued to Zoelss, G. et al.; 335,465 to Zoelss, G.; 335,464 to Zoelss, G.; and 335,467 to Zoelss, G. The pathways for preparation of celiprolol according to these patents are as illustrated:

11

35

5

1 sc

50

preparations containing celiprolol hydrochloride incorporated in a suitable carrier is applied to the glaucomatous eyes to relieve intraocular pressure. It is to be understood that both the racemic a levorotatory forms of celiprolol hydrochloride are contemplated for use in the present invention, as well as other pharmaceutically acceptable salts of celiprolol, such as maleate, succinate and the like in the range of about 0.01 to about 5% w/v.

The inactive carriers for the active compounds used ı in the formulations of the present invention include water and ointment bases, such as mineral oil in the range of about 2 to about 10% w/v and white petrolatum in the range of about 90 to 98% w/v. In preparing the formulations of the present 5 invention, the active compound is solubilized in the carrier. For solubilizing the active compound a co-solvent, in addition to the carrier, may be used. Such co-solvents include glycerin polyethylene glycol fatty acid esters in the range of 1 to 10% w/v, propylene glycol in the range of 1 to 10 10% w/v, polyethylene glycol in the range of 1 to 15% w/v, polysorbate 20, 60 and 80 in the range of 0.01 to 0.2% w/vand Pluronic F-68 in the range of 0.01 to 2% w/v and mixtures thereof.

To prevent irritation to the eye the isotonicity of the preparation should be in the range of 270 to 330 milliosmoles. Sodium chloride in the range of 0.9 ± 0.1% w/v may be used, if necessary, to adjust the isotonicity.

The ophthalmic preparations of the present invention will have a pH of about 6 to 9 preferably in the range of 7-8. Buffers that may be used to obtain said pH range include alkali metal or alkaline metal earth carbonates, bicarbonates, borates, citrates and tris buffers. More specifically such buffers include 0.01 to 0.2 molar concentrations of boric acid-sodium borate, phosphate buffers, boric acid-sodium bicarbonate, boric acid-sodium citrate, citric acid-sodium phosphate, tris(hydroxymethyl) amino methane-maleic acid and tris(hydroxymethyl) amino methane-HCl.

Other ingredients which may be desirable to use in the ophthalmic preparations of the present invention include viscosity builder agents, preservatives and stabilizers.

- Examples of these, which may be incorporated into the preparations during the process or after the active compound is solubilized, include the following:
- a. preservatives in the range of about 0.001 to about 1.0% w/v;
  - b. stabilizers in the range of about 0.01 to about 5.0% w/v; and viscosity builders or viscosity agents in the range of about 0.01 to about 2.0% w/v.

More specific examples include:

10	<u>Preservatives</u>	Range % w/v
	Benzalkonium Chloride Disodium Ethylenediamine Tetra- acetate	0.004 - 0.02 0.01 - 0.20
15	Thimerosal Chlorobutanol Phenylmercuric nitrate of the last care.	0.001 - 0.01 0.5 - 1.0 -= 0.002 - 0.02
	Methyl Paraben Propyl Paraben Phenylethyl alcohol Phenyl mercuric borate	0.002 - 0.02 0.03 - 0.20 0.01 - 0.05 0.25 - 0.75 0.002 - 0.02
20	Stabilizers	Range % w/v
25	Sodium Bisulfite Sodium Thiosulfite Cysteine Acetyl cysteine B-cyclodextrin Dextran Thiourea Thiosorbitol Monothioglyceryl disodium EDTA Dioctyl Sodium Sulfosuccinate	up to 0.5% up to 0.5% up to 3% up to 3% up to 5% up to 3%
30	Viscosity Agents	Range % w/v
•	Polyvinylpyrrolidone Polyvinyl alcohol Methyl cellulose Hydroxypropyl Methylcellulose Hydroxyethyl cellulose	$\begin{array}{ccccc} 0.5 & -2.0\% \\ 0.5 & -2.0\% \\ 0.1 & -1.0\% \\ 0.10 & -1.0\% $
35	Carboxymethyl cellulose Sodium Carboxymethyl cellulose Hydroxypropylcellulose	0.10 - 1.0% 0.10 - 1.0% 0.01 - 1.0% 0.01 - 1.0%

In general, the preparations of the present invention may be made and manufactured as illustrated herein;

- a., The active drug is dissolved in the aqueous vehicle or thoroughly dispersed in the ointment vehicle by adequate stirring;
- b., After dissolution or dispersion of the active drug additional ingredients, such as preservatives, buffer salts, stabilizers and viscosity agents are added and dissolved by further stirring. Sodium chloride is then added, if required, to adjust isotonicity, and the solution is brought to final volume;
- c., The product is then sterilized by filtration through a 0.22 micron membrane or alternatively autoclaved at 121-123°C, or by a combination of both methods;
- d., The sterile solution is filled into sterile containers and sealed.

The following examples illustrate the present invention without, however, limiting the same thereto.

20	EXAMPLE A	% w/v
	Celiprolol Benzalkonium Chloride DiSodium EDTA Purified Water, Q.S.	0.50 0.01 0.10 100.00
<b>25</b> .	EXAMPLE B	2 w/v
	Celiprolol Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate, Q.S. to pH 7.4	0.50 0.01 0.10 0.215
30	Purified Water, Q.S.	100.00

1

5

ľ

	EXAMPLE C	% w/v
5	Celiprolol Thimerosal Tris (Tribydroxyl methyl amino methane	0.125 0.002 0.12
	Maleic Acid, Q.S. to pH 8.3 Purified Water, Q.S.	100.00
10	EXAMPLE D	% w/v
	Celiprolol Benzalkonium Chloride Boric Acid	1.00 0.01 . 1.115
<b>1</b> 5	Sodium Borate, Q.S. to pH 7.4 Polyvinyl Alcohol Purified Water, Q.S.	1.4
	EXAMPLE E	: · 2 w/v
20	Celiprolol Chlorobutanol Boric Acid Sodium Thiosulfate Sodium Bicarbonate, Q.S. to pH 7.0	3.00 0.50 0.05 0.30
	Purified Water, Q.S.	100.00
25	EXAMPLE F	<u>% w/v</u>
	Celiprolol Phenyl Mercuric Acetate Methyl Paraben Propyl Paraben Polyvinyl Pyrrolidone	5.00 0.02 0.05 0.01 2.00
30	Polysorbate 80 Purified Water, Q.S.	0.05 100.00

٩.
•

Celiprolol Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate, Q.S. to pH 6.0 Polyvinyl Pyrrolidone Hydroxyethyl Cellulose Purified Water, Q.S.  Celiprolol Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate to adjust pH to 7.4 Polyethylene Glycol 300 Glycerin Sodium Thiosulfate Purified Water, Q.S.  Celiprolol Metbyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  Celiprolol Chlorobutanol Polyethylene Glycol 400			
Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate, Q.S. to pH 6.0 Polyvinyl Pyrrolidone Hydroxyethyl Cellulose Purified Water, Q.S.  Celiprolol Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate to adjust pH to 7.4 Polyethylene Glycol 300 Glycerin Sodium Thiosulfate Purified Water, Q.S.  Celiprolol Methyl Paraben Propyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  Celiprolol Chlorobutanol Polyethylene Glycol 400		EXAMPLE G	2 w/v
Polyvinyl Pyrrolidone Hydroxyethyl Cellulose Purified Water, Q.S. 100.0  EXAMPLE H	5	Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate, O.S. to pH 6.0	0.50 0.02 0.10 0.05
Celiprolol Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate to adjust pH to 7.4 Polyethylene Glycol 300 Glycerin Sodium Thiosulfate Purified Water, Q.S.  Celiprolol Methyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  EXAMPLE J  Celiprolol Chlorobutanol Polyethylene Glycol 400		Polyvinyl Pyrrolidone Hydroxyethyl Cellulose	1.50 0.20 100.00
Celiprolol Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate to adjust pH to 7.4 Polyethylene Glycol 300 Glycerin Sodium Thiosulfate Purified Water, Q.S.  Celiprolol Methyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  Celiprolol Chlorobutanol Polyethylene Glycol 400	10		
Benzalkonium Chloride DiSodium EDTA Boric Acid Sodium Borate to adjust pH to 7.4 Polyethylene Glycol 300 Glycerin Sodium Thiosulfate Purified Water, Q.S.   Celiprolol Methyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  Celiprolol Chlorobutanol Polyethylene Glycol 400		EXAMPLE H	. 2 w/v
Celiprolol White Petrolatum, Q.S.  EXAMPLE J  EXAMPLE J  Celiprolol Coliprolol Coliprolo	15	Benzalkonium Chloride DiSodium EDTA Boric Acid	0.50 0.005 0.05 0.215
Celiprolol Methyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  EXAMPLE J  Celiprolol Chlorobutanol Polyethylene Glycol 400		Polyethylene Glycol 300 Glycerin Sodium Thiosulfate	5.0 1.0 0.5 100.00
Methyl Paraben Propyl Paraben Mineral Oil White Petrolatum, Q.S.  EXAMPLE J  Celiprolol Chlorobutanol Polyethylene Glycol 400	20	EXAMPLE I	<u>% w/v</u>
Celiprolol Chlorobutanol Polyethylene Glycol 400	25	Methyl Paraben Propyl Paraben Mineral Oil	1.0 0.05 0.01 5.0 100.00
Celiprolol Chlorobutanol Polyethylene Glycol 400		EXAMPLE J	2 w/v
20 morre leriotatum. O.S. jun v	30	Chlorobutanol	0.5 0.5 4.0 100.00

Illustrative of the benefits obtained in accordance with the present invention the studies described in the Examples following, were conducted.

#### EXAMPLE 1

This example shows that celiprolol decreases intraocular pressure in dogs and that the decrease is dose-dependent.

### Animals, Testing Procedure and Apparatus

Mongrel dogs of either sex weighing between 9 and 16 kg were anesthetized with sodium pentobarbital, 35 mg/kg, i.v. (Ganes Chemical, Pennsville, N.J.). The animals then were intubated with an endotracheal tube (Rusch, size 8-9F, Artistic Surgical, New York, N.Y.) and allowed to breathe spontaneously.

Pulsatile arterial pressure was monitored using a 15 polyethylene catheter (PE 240, Clay-Adams, Parsippany, N.J.) inserted into the right femoral artery and its tip advanced until a distinct dicrotic notch was observed on the arterial pressure tracing. The catheter was connected to a pressure transducer (P23 ID, Statham, Oxnard, CA) and a D.C. driver 20 amplifier (Model 7D; Grass Instruments, Quincy, MA) via a low leve D.C. preamplifier (Model 7P1, Grass Instruments). Mean arterial pressure was determined electronically by damping the pulsatile arterial pressure signal. Heart rate was recorded from the output of a tachometer (Model 7PA, Grass 25 Instruments) triggered by the R wave of a Lead II electrocardiogram (EKG -Tachograph Preamplifier, Model 7P4, Grass Instruments). The outputs of arterial pressure and EKG -Tachometer Preamplifier were recorded continuously on an oscillograph (Model 7D, Grass Instruments). Heart rate was 30 measured manually from the tachometer output or calculated from the EKG preamplifier output. Mean arterial pressure was measured manually from the oscillograph tracings.

Intraocular pressure was measured using a pneumatonometer (Model 30R, Digilab, Cambridge, MA).

Upon completion of all surgical procedure the animals were allowed to stabilize for 15-30 minutes before pretreatment baseline measurements were taken. PROTOCOL

Eight mongrel dogs were divided into two groups of four dogs each: the control group and the test group. Two pretreatment baseline measurements of intraocular pressure, mean arterial pressure and heart rate were recorded at -15 and 0 minutes. The means values of these two readings were used as the pretreatment (0 time) values. The control group was then administered saline topically in a volume of  $5\,\mu\mathrm{l}$ instilled into the left eye of each dog at hourly intervals. The test group was administered celiprolol topically in a volume to 50 Hl instilled into the left eye of each dog, in ascending concentrations (0.03%, 0.06%, 0.125%, 0.5%), at hourly intervals. Intraocular pressure, mean arterial pressure and heart rate were recorded at 15 minute intervals for 60 minutes post drug or vehicle (saline) administration for each concentration of celiprolol administered and each administration of saline.

#### Drug Preparation

Celiprolol (RHC 5320-A Lot 2) was dissolved and diluted in normal saline (0.9%, Abbott Laboratories, North Chicago, IL).

The control group received normal saline.

#### Data Analysis

The absolute change and the percent change from the pretreatment (0 time) values for intraocular pressure, mean arterial pressure and heart rate were expressed as the mean to S.D. The significance of the difference between the vehicle

5

10

15

20

control group and the celiprolol test group were evaluated using a "t" test for grouped data (as described in SAS T TEST Release 79.5, SAS Institute, Gary, NC 1982). Differences were considered significant if P 0.05. In addition, the maximum percent intraocular pressure change for each concentration of celiprolol administered was expressed as the

Result

mean  $\pm$  1 S.D.

Test results were shown in Tables I, II & III.

10

15

20

25

30

48° 41

-44p

145 b

142

-28 -28

4 7 71

-33 +15

-43p

-37<sup>b</sup>

4 5t 21

-33<sup>b</sup> -37<sup>b</sup> +11 +8

-33<sub>b</sub>

-15 -16 -28

7 F

1 Change

-15<sup>b</sup>

14 -14

17. FP

14 L

9 7<sub>1</sub>

4,7°F1

1+75

479

74.

\$ T ₹₁

-10<sub>p</sub>

-10<sup>b</sup> -11<sup>b</sup>

-10p

T 71

ን <del>ፒ</del>Ι ა. <u>‡</u>1

7 71

Change on Hg

2 71

Saline (50 pl of 0.9% NaCl Solution)

			300	4 ‡!	271
1		0.5%	285	۳ <del>۲</del> ۱	-25 +6
		ó	270	7 <del>7</del> 1	+13
F			255	2 11	-23 +16
5			240	<b>ግ</b> ∓ι	1+3
	DOGS	0.25%	225	. ሷl	-24 +5
	0821138	0.0	210	9 71	-53 +4
10	ANEST		195	\$7 <del>7</del> 1	-22  +8
	NI (ac		180	~ <del>~</del> 1	-21 +16
	RE (IC	125%	165	Ÿ <b>‡</b> I	, 5 <sup>7</sup>
15	PRESSU	0.125%	150	4 5 <sup>1</sup>	77 PJ
	CULAR	-	135	771	-13 -11 -11
	INTRAC		120	4 41	-18 -11
20	OL ON	0.062	103	4 4	-17
	TABLE I		90	7 F1	B- (1)
	30 40		75	7 71	77
	TONS		90	7 7 <sub>1</sub>	7 %1
25	TRAT	2.032	£5.	771	14 41
	DNCEN	0	30	7 PI	4 17
	ING C		13	041	2 -6 ±10 ±13
30	TABLE I THE BPPECT OF ASCENDING CONCENTRATIONS OF CELIPROLOL ON INTRAOCULAR PRESSURE (IOP) IN ANESTHETIZED DOGS	Baseline Dose Time	Post- trestment	Change vo Hg	% Change
	THE BFFEC	Baseline <sup>a</sup> Pre-	treatment (mm Hg)	21 +2	
			z	•	

35

All values are the mean  $\pm$  1 5.D. bp<0.05

					_			
				ğ	- <del>-</del> +11	φ <sup>4</sup> 1	4.20	771
1				285	5 11 <del>1</del>	3 +10	9 + 1	1+12
			0.52	270	177	0 +11	-4 +11	7 %
			•	255	<b>~</b> ¥ı	9 <del>7</del> 1	-5 +16	+1+
5				240	~ 71	r <del>1</del> 1	-5 +16	-5 +14 144
	9068		-	225	9 41	9 41	4 1	r 41
	TIZED		0.3%	210	9 +10	~ <del>\$</del> 1	£ <del>4</del> )	7 7 <sub>1</sub>
10	NES THE			195	12 +14	# <del>[</del> ]	-4 +10	4 <b>L</b> !
	4 H			180	177	6 +15	11	10
•	E CKAP			165	8 +16	7	7 <del>1</del>	77
15	RESSIT		0.1%	150	11	10	791	" 14 17
	RIAL			35	6 1+15	5 113	-7 +10	<b>ም</b> ምι
	X ARTE			120	10	8 +10	<u>ر. 4</u> ا	9 (+)
20	II NO NO		ж	105	o <del> </del> 11	e 41	9-19-1	4 1.5 to
20	TABLE II		190.0	06	ا4 ه	8 41	4B P 141	罗乳
	Kal Lai			22	4+18	8 +16	771	<sub>የ</sub> ታ ኢነ
	9		<u> </u>	99	e ∓1	~ \$1	77 军(	7 4
25	SNOTE		н	. <b>:</b> 3	4 111	170	T <b>T</b> 1	7 4
_	4 OTM3:		0.03%	2	6 ±15	5 <u>+13</u>	2 +15	2 +11
	מט טאַ			15	æ <b>°</b> †1	~\$1	110	° <del>4</del> 1
30	TABLE II.  TABLE II.	1	Dose Time	Post- trestment	Change (um Hg)	1 Change	Change (mm 11g)	7 Change
	Loddad Ant	100 010		treatment N (mm Hg)	113		126 <u>+</u> 15	
25				×	¥ -		4	
35				Trestment	Saline (50 µl of 0.9% 4 NaCl Splution)		Celiprolol HCl (RHC 5320-A-2 50 µl)	

All values are the mean  $\pm$  1 S.D. bp<0.05

			!	90
ı			н	185
			0,52	270
		•		255
5				240
			H	225
		D DOGS	0.3%	195 210 225
10		ETIZE		195
		ANEST		180
		(S)	, e	165
15		VTE (BF	0.13	150 165
		ART R		135
		ON RE		120
20	TABLE III	PROLO	)6Z	75 90 105
	TAB	F CELL	0.062	8
		O SNO		
		TRATI		30 45 60
25		ONCEN	0.03%	A.
		ING	0	30
		SCEND	L	23
30		THE EFFECT OF ASCENDING CONCENTRATIONS OF CELIPROLOL ON REART RATE (BPN) IN ANESTHETIZED DECS	Dose Time Minutes Post-	trestment
		THE	Baseline Dose Time Fre- Minutes	N (om Hg)
35	•			

	300	1+31	1757	+35	7 2
н	:83	-27	-18 +10	-18 +29	-13
0.52	270	-30 +29	-20 +19	8. ±1	\$- Ef.
	255	-23	-15 +10	-11 +25	-7 +18
	240	9-1450	177	-17	-12 +13
7	225	175	-17 +15	117	-12 +8
0.3%	210	-37	-21 -15	77	, <del>, ,</del> ,
	195	-24 +35	-16 +23	133	- <del>1</del>
	180	-23 +35	-15 -15	971	5 +20
1%	165	-17 +28	11-	P 🛱	4 21
0.1%	150	17 F	-16 +21	2. E	1, <del>1,</del> 1,
	135	-24 +35	-16 123	122	11.
	120	6+1	4 4	-10  +16	-7 +12
25	105	178	ē. ±1	6 +1	-7 +20
0.052	96	-18 +21	-12 +14	-6 +15	2. <u>†</u>
	75	-23 +17	감다	177	4 <u>4</u>
	9	771	77	م <del>با</del> با	7 71
,03£	4.5	-17	79	1412	741
6.0	30	477	9 41	4 41	771
	2.5	9 F1	9 4	0 +1	o 71
Dose Time Minutes	treatment	Change (bpm)	Change	Change (bpa)	% Change
Baseline <sup>6</sup> Pre-	(nm Hg)	153		135 +14	•
	×	4		4	
	Treatment	Saline (50 µl of 0.9% NeCl Soluțion)	٠	Celiprolol HCL (RHC 5320-A-2) 50 µl)	
		~			

All values as the mean  $\pm$  1 S.D. b P < 0.05

Table I shows that celiprolol caused an immediate decrease in intraocular pressure which persisted for the duration of the experiment. The magnitude of the decrease became significantly different (P<0.05) from that of the control group at 75 minutes. The data also indicate that celiprolol causes a dose-dependent decrease in intraocular pressure which reaches a maximum at 0.25% concentration.

Table II shows that the celiprolol group exhibited a small decrease in mean arterial pressure which became significantly different from that of the control at a concentration of 0.06%. Since this decrease in arterial pressure was not dose-dependent, it may be a chance occurence and not pharmacologically important.

Table III shows that celiprolol had no effect on heart rate at any of the concentrations tested.

20

10

25

30

#### EXAMPLE II

This example compares the activity of celiprolol and timolol in dogs on intraocular pressure, systemic arterial pressure and heart rate.

### Animals, Testing Procedure and Apparatus

Same as described in Example I.

#### Drug Preparation

Celiprolol and saline were used as shown in Example I. Timolol (TIMOPTIC , 0.5%, Merck Sharp and Dohme, West Point, PA) was used.

#### PROTOCOL

1

5

10

Mongrel dogs were screened for intraocular pressure and only dogs that had intraocular pressures between 23 mmHg and 28 mmHg were chosen in this study.

Twelve mongrel dogs which met the above-described criterion were subdivided into three groups of four dogs each. The control group had a mean intraocular pressure of 25.3 ± 0.6 mmHg; the celiprolol, or test, group had a mean intraocular pressure of 25.6 ± 0.6 mmHg; and the timolol group, with which the celiprolol group was compared, had a mean intraocular pressure of 25.2 ± 1.6 mmHg.

Two pretreatment baseline measurements of intraocular pressure, mean arterial pressure and heart rate were recorded at -15 and 0 minutes for each groups. The mean values of these two readings were used as pretreatment (0 time) values 0.5% celiprolol in a volume of 50 $\mu$ 1 was then administered into the left of each dog in the test or celiprolol group; 0.5% timolol in a volume of 50 $\mu$ 1 was administered into the left eye of each dog in the timolol group; and 50 $\mu$ 1 of saline was administered into the left eye of each dog in the control group. Intraocular pressure, mean arterial pressure and heart rate measurements were made at various intervals for five hours respectively for post drug or saline administration.

#### Date Analysis

Same as in Example 1.

#### Results

٠1

Test results are shown in Tables IV, V and VI Table

IV shows that at equal doses both celiprolol and timolol were effective in decreasing intraocular pressure in anesthetized dogs. Celiprolol, however, caused a significant change from control of the one hour post drug administration which persisted for the duration of the study, while timolol attained a significant difference from the control only after 180 minutes post drug administration, which from thereon also persisted for the duration of the experiment.

Table V shows that neither celiprolol nor timolol had any significant effect on means arterial pressure.

15 effect on heart rate whereas timolol at least for the first two hours has a tendency to decrease heart rate.

20

25

30

1	
5	
10	
15	
20	
25	

TABLE IV

- Dogs								
THETIZED		300	771	9 +	977	-41 <b>6</b> -41 <b>6</b>	<sub>ቅ</sub> ትነ	-30 <sub>b</sub>
IN ANES!		270	<b>→</b> 71	43	1 7 71	-38 b	5. th	-27 <sup>b</sup>
(10P)	٠	240	- 41	-4 +19	14 27	-36 p	17 <sup>9</sup>	416-31
RESSURE	ac)	210	0 71	-1 +15	<b>₹</b>	-37b	P 71	-30p
CULAR P	Time (Minutes Posttrestment)	180	o \$1	17 -1	48 21	-31 <sub>b</sub>	<b>4</b> 51	-31 <sub>6</sub>
INTRAO	es Post	120	- ፕ೪	- st.	<b>4</b> ₹1	-33°	ዋኍ <sub></sub> ነ	-32 +20
NO 7070	(Minut	120	.7 Pj	9 취	. \$ 71	-23°b	741	-25 +16
TIM	Tine	90	0 +1	7 41	45.2 <sup>1</sup>	-21 <sub>b</sub>	441	-22 +15
TOT AN	;	90	o 71	77,	771	-53 <sub>p</sub>	771	17 F1
CELIPRO	:	45	° 71	1 +16	7 71	110	ኔታl	-20 +20
TERED	ć	Ş	7 \$1	4 27	T 71	11-51	741	다.
ADMINIS	;	3	74,	ς <del>1</del> 1	7 PI	감함	771	77
THE EFFECT OF TOPICALLY ADMINISTERED CELLPROLOL AND TIMOLOL ON INTRAOCULAR PRESSURE (10P) IN ANESTHETIZED DOCS			Change (cm Hg)	X Change	Change (mm Hg)	Change	Change (rm Hg)	X Change
FECT OF 1	Baseline <sup>8</sup> Pre- treatment (mm Ho)	/9	14 25		17.3		. 1+2	
3112	*	i	4 ution)		1 4 luțion	-	<b>5</b>	
	Trestment		Saline 50 pl (0.9% NaCl Solution)	i	Celfproiol HG1 (RIC 5320-A) 50 µl 0.5% Solution		Timolol Maleate (Timoptic) 50 µl of 0.5 Solution	q

All values are expressed as the mean  $\pm$  1 S.D. bp < 0,05

35

30	20		25			20			15		_~	10		5		1
								27	TABLE V	3	F03 748	PRTAL P	RESSURE	(WAP)	IN ANESTHE	TZED DOGS
		THE BFFECT	OF TOPIC	VILY AL	MINIS	TERED (	ELIPRE	NOT WE	DIMORG	200	EAST ARE	To the same of the			THE BFFECT OF TOPICALLY ADMINISTERED CELIPROLOL AND TIMOLOL ON NEAR ANTENDED.	
	Ā	Baseline					7	K 98 (A	Time (Minutes Posttreatment)	osttre	atment)					
7. 4. 1. 1. 1.	y z	Pre- treatment (mm Hg)		21	R	45	09	26	120 1	150	180	210	240	270	300	
Saline 50 ul 0.9% NaCl Solution)	4	120 <del>1</del> 13	Change (mm Hg)	4 FI	n #l	n¥l ·	n <del>9</del> 1	4 <del>1</del> 1	+11 +	2 +10	n <del>T</del> 1	5 +13	2 <del>11</del> 2	o #l	v <del>4</del> 1	
			% Obange	n #1	n <del>1</del> 1	n #i	1 <del>7</del> 2	2 <del>4</del> 1	_	~ <del>1</del> 1	 •≠*¥I	4 +12	1410	o <b>^</b> 1 1	<b>4₽</b> 1 Ϊ	
Celiprolol HG1	4	611 8+	Change (mm 11g)	7	- FI	4 #1	n #1	n #1	17 19	0 ±12	n #i	~ <del>}</del> !	n <del>1</del> 11	7 <b>2</b> 1	· ‡.	
50 µl 0.5% Solution	e	i	7 Change	n <sup>2</sup> 1	<b>4</b> ₹i	n ¥i	<b>4</b> 41	n 41	o <del>(</del> 1	6+10	en <del>(</del> )	~ <del>+</del>	~¥I	771	79)	
Timolol Maleate (Timopeic)	4	108	Change (rm 11g)	7 <del>7</del> 1	7 %	77		147	97 +1	<b>₹</b>	T \$1	<b>다 </b> 뛰	<i>ል ኢ</i> !	7 <b>4</b> 1	<b>ጉ</b> ፟	
Solution			% Change	7 41	741	741	<b>4</b> 41	e 21	+23	4.15 5.15	7 %	<b>다</b> %	471	7 <del>(</del> )	7 %	
All values are expressed as the bp<0.05	presse	id as the m	mean + 1 S,D,	ė				•	: -							

1			2000									
5			ETIZED		300	-12 +18	-10 +14	-28 +10	-22 -25	-23 +26	-27 +22	
			ANESTH		270	-13 +30	-10 +21	-18 -21	-14 +16	14.8	7 F1	
			BMP) IN		240	131	6- 43	177 1733 1733	-14 +17	-26 +32	-18 +29	
10			RATE (	atment)	210	-5	£ 51	77	P 71	-23 +37	-15	
			N HEART	Tine (Minutes Posttrestment)	180	-130 -130 -130	17 57	-11 +16	6- 21-	-28 +33	-19 +29	
			HOLOL O	Inutes	150	6 <del>,</del> 49	1년8	-21 +20	-17 -18	. <sup>17</sup> <sup>13</sup> .	-17 -13 -13 -13	
15	_		AND TI	Yne (M	120	8 110 110	ا‡ م	-5 118	₹. •	141	-10	
	•	TABLE VI	ROLOL	F-	90	o Ŧ1	411	44 111	الة با	-25 +25	-18 +21	•
		TABL	CELIP		9	07+	179	2 1	4-11	-28 +18	-22 ±13	
20			STERED		45	3	3 +12	+18	δ <u>‡</u> !	-25 <sup>b</sup> +17	-19 <sup>b</sup>	
			ADMIN		8	5 +20	3	-10 +10	T \$1	-19 +11	17 PI	
			TCALLY		15	6 ±11	411	5 \$1	7 71	111	9 Pi	ė,
25			THE BFFECT OF TOPICALLY ADMINISTERED CELIPROLOL AND TIMOLOL ON HEART RATE (BMP) IN ANESTHETIZED DOGS			Change (bpp)	Z Change	Change (bpm)	7 Change	Change (hps):	Z Change	the mean + 1 S,D,
			THE KFF	Baseline <sup>a</sup> Pre-	(ma Kg)	122		137		119 <del>1</del> 22		
30					2	4 lucton		1 4 lution		ce 4 Solution		e expresse
35					Treatment	Saline 50 µl 0.9% NaCl Solution		Celiprolai HC1 (RHC 5320-A) 50 µl 0.5% Solution		Timolal Maleate 4 (Timoptic) 50 ul of 0.3% Solution		All values are expressed as

In asthmatic patients airway constriction has been reported as the result of the use of certain -blockers. The present invention, utilizing celiprolol, provides for the treatment of glaucoma without having the deleterious side effect of airway constriction associated with certain \$\mathcal{B}\$-blockers, such as atenolol, metoprolol, propanolol and timolol. Example III describes the comparative study conducted on celiprolol with three other beta blocking agents for effects on bronchomotor tone in mechanically ventilated cats infused with serotonin.

#### EXAMPLE III

#### METHOD

1

Male cats (3-5 kg) were anesthetized with pentobarbital sodium, 35 mg/kg i.p. Maintenance anesthetic was administered intravenously as needed. The trachea of each cat was cannulated and pneumotachograph placed on-line for monitoring air flow. A differential pressure tranducer placed between the tracheal cannula and a cannula in the pleural cavity was used to monitor transpulmonary pressure. Electronic signals proportional to air flow and 10 transpulmonary pressure were converted by an on-line analog computer to values of pulmonary (airway) resistance, RAW' for each breath. Cannulae were inserted into the right femoral artery and both femoral veins. The arterial cannula was used for monitoring blood pressure and heart rate. 15 addition, it allowed for withdrawal of arterial blood samples for assessment of PO2, PCO2 and pH, as a means of confirming the adequacy of ventilation. The animal was paralyzed with gallamine triethiodide (Flaxedil, 20 mg i.v.) and mechanically ventilated. 20

In order to test for bronchodilator activity, it was necessary to increase the normally low bronchomotor tone of the cat. This increase was induced by a constant i.v. infusion of serotonin (5-HT), approximately 20 /kg/kg/min. During this steady state bronchoconstriction, each animal received three or four increasing bolus doses of a single beta blocker, either atenolol (1-10 mg/kg/i.v.), celiprolol hydrochloride (1-10 mg/kg i.v. or 1-100 mg/kg i.d.), metoprolol tartrate (1-10 mg/kg i.v.) or timolol maleate (0.03-3 mg/kg i.v.). Doses are expressed as the free bases. All drugs were dissolved in 0.9% saline, except timolol, which was used in the form of Timoptic (1-10 mg/kg).

- solution suitably diluted with saline. In exp riments in which celiprolol was given introduodenally, needle-tipped cannulae were surgically inserted into the duodena of cats fasted for 16 hours.
- Drug-induced changes in bronchomotor tone, averaged over 6-second intervals, were calculated as percent changes in R<sub>AW</sub> from the steady state values established by serotonin infusion. Mean ± standard deviation were calculated for experiments in which n = 3. Data for one celiprolol dose, 10 mg/kg i.d., were calculated using n = 2. Statistical analysis was carried out with the t-test for paired data.

#### RESULTS

The results are shown in Table VII wherein R<sub>AW</sub> = 15 % change in airway resistance; HR = heart rate - breaths/.
minute; i.v. = intravenous; i.d. = intraduodenal.

#### TABLE VII

20	COMPOUND	mg/kg i.v.	HR	RAW
	ATENOLOL	1 3 10	-8 <u>+</u> 5 -6 <u>+</u> 3 -6 <u>+</u> 2	23+12 14 <del>1</del> 5 48 <u>+</u> 34
25	METOPROLOL	1 3 10	-17+11 -25∓19 -94 <u>∓</u> 86	50 <u>+23</u> 43 <u>+</u> 21 105 <u>+</u> 37
30	TIMOLOL	0.3	-8+8 -8 <u>+</u> 1	11 <u>+</u> 9 18 <u>+</u> 51
	CELIPROLOL	1 3 10	-1+10 -9+4 -27+7	-62+10 -65+10 -70+14

As shown in the table, intravenous administration of atenolol, metoprolol and timolol caused dose-dependent bronchoconstriction, i.e., increased R<sub>AW</sub>. Bronchoconstriction has increased to 48% with atenolol, to 105% with metoprolol and to 49% with timolol. In contrast, celiprolol produced bronchodilation, i.e., R<sub>AW</sub> decreased to -70%.

The duration of the broncodilator effect of celiprolol was about 14 minutes at 1 and 3 mg/kg. The effect of 10 mg/kg lasted for more than 40 minutes.

EXAMPLE IV

The procedure in this example generally follows the procedure used in Example III, except that the test compounds of celiprolol and timolol were administered topically as follows. 50 ½ 1 of test solution was instilled into the eyes of anesthetized cats whose airway resistance, R<sub>AW</sub>, had been increased by infusion of serotonin. Three doses of timolol (5 mg/ml) further increased R<sub>AW</sub> by 20 to 35%. Similar doses of celiprolol decreased R<sub>AW</sub> by 21 to 23%.

0109561

ī

#### EXAMPLE V

This procedure in this sample generally follows the procedure used in Example III, except that celiprolol was administered to cats intraduodenally. Tested in the range of 10-100 mg/kg, celiprolol; reduced  $R_{\rm AW}$  by 29 to 41%.

Having described the invention, those skilled in the art will know modifications within the spirit thereof, and the invention is to be limited only within the scope of the appended claims.

10

15

20

25

30

### WHAT IS CLAIMED IS:

1

5

- 1. An ophthalmic preparation for the treatment of glaucoma comprising an effective amount of a c liprolol salt in a pharmaceutically acceptable ophthalmic carrier for lowering intraocular pressure.
- 2. The ophthalmic preparation of Claim 1 wherein the celiprolol salt is present from about 0.1 to about 5.0% w/v concentrations in said pharmaceutically acceptable carrier.
- 3. The ophthalmic preparation of Claim 1 or 2 wherein from about 0.03 to about 0.5% w/v of the celiprolol salt is present in a pharmaceutically acceptable carrier.
  - 4. The ophthalmic preparation of Claim 2 or 3 wherein said pharmaceutically acceptable carrier comprises in % w/v:

15 boric acid-sodium borate 0.03 benzalconium chloride 0.01 disodium ethylenediamine 0.1 tetraacetate sodium thiosulfite 0.3 20 1.5 polyvinylpyrrolidone 0.1 hydroxyethyl cellulose 0.01 polysorbate 80 water QS to 100, and having an isotonicity range of 270 to 330 milliosmoles. 25

- 5. The ophthalmic preparation of any of Claims 1 to 4 wherein said pharmaceutically acceptable carrier is a water base or an ointment base.
- 6. The ophthalmic preparation of any of Claims 1

  to 5 wherein the isotonicity of said preparation is within the range of 270 to 330 milliosmoles.

- 7. The ophthalmic preparation of any of Claims 1 to 6 wherein the pH of said preparation is in the range of 6 to 9.0.
- 8. The ophthalmic preparation of any of Claims 1 to 7 wherein said celiprolol salt is celiprolol hydrochloride.
  - 9. The ophthalmic preparation of any of Claims 1 to 8 further comprising a buffer, a tonicity agent, a preservative, a stabilizer, a co-solvent and a viscosity agent.
  - 10. An ophthalmic preparation for the treatment of glaucoma according to Claim 1 comprising:
    - a. celiprolol hydrochloride 0.03-2.0 w/v;
- b. a co-solvent selected from the group consisting of glycerin, propylene glycol, polyethylene glycol, polysorbates 20, 60 and 80 or Pluronic F-68;
  - c. a viscosity agent selected from the group consisting of polyvinylpyrrolidone, polyvinyl alcohol, methyl cellulose, hydroxypropyl methylcellulose, hydroxyethyl cellulose and carboxymethylcellulose; sodium carboxymethylcellulose or hydroxypropylcellulose;
  - d. a stabilizer selected from the group consisting of sodium bisulfite, sodium thiosulfite, cysteine, acetyl cysteine, B -cyclodextrin, dextran and thiourea, thio-sorbitol, morothioglycerol, sodium EDTA, or sodium sulfosuccinate:
- e. a 0.01 to 0.20 molar buffer selected from the group consisting of boric acid-sodium borate, phosphate buffer, boric acid-sodium bicarbonate, boric acid-sodium citrate, citric acid-sodium phosphate, tri(hydroxymethyl) amino methane-maleic acid and tris(hydroxymethyl) amino methane-HCl;

10

20

- f. a preservative selected from the group consisting of benzalkonium chloride, disodium ethylene-diamine tetracetate, thimerosal, chlorobutanol, phenylmercuric nitrate, phenylmercuric acetate, methyl paraben, propyl paraben, phenyl mercuric borate or phenylethyl alcohol; and
  - g. water Q.S. to 100%.
  - 11. A process of preparing an opthalmic preparation for lowering intraocular pressure comprising:
- a. dissolving or dispersing an effective amount of celiprolol salt in a pharmaceutically acceptable carrier;
  - b. adding buffer salts, stabilizers or viscosity agents;
  - c. further optionally adding sodium chloride to adjust to the required isotonicity;
  - d. adjusting to the final volume with purified water.
  - 12. The process according to Claim 1 wherein the preparation is further sterilized by filtration or autoclaved, or both.

30

15



EPO Form 1503 03 67

## **EUROPEAN SEARCH REPORT**

Application number

EP 83 11 0477

	Citation of document	ISIDERED TO BE RELEVA		
Category	of to	with Indication, where appropriate, elevant passages	Relevant to Claim	CLASSIFICATION OF THE APPLICATION (Int. CI. 3)
D,X	DE-A-2 458 624 * Claims; exam	(LENTIA) ple 1 *	1-12	A 61 K 31/1
	-			
	•			
	·		-	TECHNICAL FIELDS 8EARCHED (Int. Ci. 3)
				A 61 K 31/00
	The present search report has b	een drawn up for all claims	-	
	Place of search THE HAGUE	Date of completion of the search 02-02-1984	GAUTIER	Examiner R.H.A.
: partic : partic docum : techno	CATEGORY OF CITED DOCU ularly relevant if taken alone ularly relevant if combined with nent of the same category ological background	E : earlier pate	rinciple underlying ant document, but ping date cited in the applicated for other reasons.	the invention
	ritten disclosure rediate document			mily, corresponding